



Rodent diversity in South America: transitioning into the genomics era

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The goal of this perspective is to propose a concerted effort for the study of the diversity of South American rodents to transition into the genomic era. We first review progress made in the last few decades in our understanding of the age, geographical origins, phylogenetic relationships and diversity of caviomorph and sigmodontine rodents, two major components of the South American fauna. Then, we examine their current representation in genomic/transcriptomic databases, and outline a research program to obtain a set of genomes and transcriptomes that represent major groups within each of these two lineages. We propose standards for voucher specimens and associated data and explore the role of museums and research collections in this endeavor. We envision a concerted international effort to guide and accelerate progress in the field.

Keywords: Hystricognathi, museums, phylogenomics, Rodentia, Sigmodontinae, South America

INTRODUCTION

The field of phylogenetics is undergoing the transition to phylogenomics. As sequence technology progresses, associated costs reduce, computational capacity increases, and analytical protocols improve (e.g., Chan and Ragan, 2013; Dunn et al., 2013), the transition from traditional molecular systematics based on few loci to matrices of hundreds or thousands of genomic regions becomes feasible for most organisms. This transition is now underway and its pace will quicken in the next 5 years. While the analysis of genomic scale data would not solve all phylogenetic questions, it would help address the most important caveat of molecular systematics, namely the difference between gene and species trees resulting from sorting of ancestral polymorphism and introgression and magnified by difficulties of accurately reconstructing individual gene trees. Analysis of genomic scale data reduces the uncertainty of phylogenetic inference and consequently increases the power of tree-based hypothesis testing.

South America hosts some 1300 mammalian species, representing about 23% of all living mammals (<http://www.iucnredlist.org/initiatives/mammals/analysis>). Considerable progress has been made in our understanding of the origin and evolution of the impressive diversity of South American mammals, due primarily to fossil discoveries and DNA sequence-based phylogenetic analysis (see Patterson and Costa, 2012). The transition of the field of phylogenetics and systematics into the genomic era promises new insights into the diversity of South American mammals and their functional roles in diverse ecosystems. We believe progress in this exciting area of research will be greatly accelerated if a concerted, international research program stimulates the production of a core set of high quality, large scale genomic and transcriptomic reference sequences for a select set of mammalian species. Holistic voucher specimens with

rich data and materials that are made widely available would then facilitate expansion in multiple directions in the future, including ecology, physiology, developmental biology, and public health arenas.

To outline such a research program and examine its feasibility, we focus on rodents. We specifically concentrate on caviomorph and sigmodontine rodents, two groups largely (but not exclusively) found in South America because collectively they encompass 614 (95%) of the 642 species of South American rodents (Patton et al., in press). Focusing on sigmodontines and caviomorphs allows us to provide specificity in terms of both challenges and opportunities. This exercise aims to stimulate the overall effort and may also serve as an example that can be extended to other taxonomic groups. We first provide an overview of the progress made in phylogenetics and systematics on the basis of DNA sequence data; then, we examine how a genome-level program might help resolve major pending issues and, more generally, facilitate advances in various fields.

CAVIOMORPH RODENTS: AGE, ORIGINS AND DIVERSITY

Caviomorphs (guinea pigs and allies, comprising 46 genera and 233 species in South America, **Table 1**) are unusual among rodents because some reach large body sizes (e.g., >50 kg in the capybara), have relatively long gestation times, and diverge from other mammals at genes that are usually conserved, such as insulin (Opazo et al., 2005) and growth hormones (Adkins et al., 2000). The phylogenetic position and geographical origin of caviomorphs has been controversial. Three decades ago, the fossil record of caviomorphs traced back to only about 22 Mya (reviewed by Reig, 1981), their monophyly relative to other hystricognath rodents (phiomorphs: African mole rats, dassies, and cane rats, and hystricids: Old World porcupines) was questioned

Table 1 | Classification and diversity (number of species and genera per family) of caviomorph rodents in South America (following Patton et al., in press), with a survey of the available taxonomic coverage for the most commonly used loci in caviomorph phylogenetic studies, and available and proposed genomic and transcriptomic coverage.

Super family	Family	Genera (species)	Genes most commonly used in phylogenetics						Available mitochondrial genomes	Available* and proposed (minimum) coverage		
			Mitochondrial		Nuclear					Genomes (NCBI accession)	Transcriptomes	
			cyt b	12 s	IRBP	vWF	TTH	GHR				
Cavioidea		9 (35)	9 (21)	9 (16)	4 (5)	4 (6)	9 (15)	9 (16)	1	1* (PRJNA73553)	2*+ 2	
	Caviidae	6 (21)	6 (16)	6 (11)	2 (2)	2 (2)	6 (10)	6 (10)	1 1	1*	2*	
		Cuniculidae	1 (2)	1 (1)	1 (1)	1 (2)	1 (2)	1 (2)	1 (2)			1
	Dasyproctidae	2 (12)	2 (4)	2 (4)	1 (1)	1 (2)	2 (3)	2 (4)			1	
	Chinchilloidea		4 (8)	4 (6)	4 (2)	2 (2)	2 (2)	4 (4)	4 (4)	1	1* (PRJNA68239)	1*+ 1
		Chinchillidae	3 (7)	3 (5)	3 (1)	1 (1)	1 (1)	3 (3)	3 (3)	1 1	1*	1*
Dinomyidae			1 (1)	1 (1)	1 (1)	1 (1)	1 (1)	1 (1)	1 (1)			1
Erethizontoidea			2 (13)	2 12	1 (4)	0 (0)	1 (1)	1 (1)	1 (1)	1	1 (PRJNA15171)	1
		Erethizontidae	2 (13)	2 (12)	1 (4)	0 (0)	1 (1)	1 (1)	1 (1)	1 1	1	1
Octodontoidea		31 (177)	24 (90)	23 (45)	10 (19)	19 (42)	11 (12)	21 (40)	6	1* (PRJNA193441)	2*+ 2	
	Abrocomidae	2 (10)	1 (2)	1 (1)	1 (2)	1 (1)	1 (1)	1 (2)			1	
		Ctenomyidae	1 (65)	1 (30)	1 (3)	1 (1)	1 (3)	1 (1)	1 (3)	1 3		1*
	Echimyidae	22 (88)	16 (49)	15 (30)	6 (14)	15 (35)	3 (3)	13 (24)	2 2		1	
		Octodontidae	6 (14)	6 (9)	6 (11)	2 (2)	2 (3)	6 (7)	6 (11)	3 3	1*	1*

The *Capromyidae* do not live in South America but should be included in phylogenomic efforts to complete all families of caviomorphs. Genome projects are identified by NCBI accession numbers; additional studies: mitochondrial genomes are reported in Tomasco and Lessa (2011) and Voloch et al. (2013); transcriptome in MacManes and Lacey (2012).

by myological and immunological analyses, and there was strong disagreement between proponents of an African connection to phiomorphs and champions of a North American origin.

Many of those controversies have subsided. Molecular data support the monophyly of caviomorph rodents and this clade is now widely thought to be sister to phiomorphs and to have originated from a single colonization from Africa (e.g., Cao et al., 1994; Frye and Hedges, 1995; Huchon and Douzery, 2001; Honeycutt et al., 2003; Rowe et al., 2010). DNA sequence analyses suggest that the divergence between caviomorphs and phiomorphs occurred 43–45 Ma, and that the early branching of caviomorphs took place 37–38 Ma (Poux et al., 2006; Voloch et al., 2013), effectively doubling the age of the group in South America.

Recent fossil findings document the presence of already diverse caviomorphs as early as 41 Ma in South America (Vucetich et al., 2010; Antoine et al., 2012; Bertrand et al., 2012).

In spite of these significant advances, many unresolved issues remain that will require considerably greater genetic and taxonomic sampling. Among them, we highlight the following:

- Higher level relationships.* The organization of living caviomorph families into four superfamilies appears to be well-supported (Upham and Patterson, 2012). In turn, Erethizontoidea and Cavioidea form a well-supported clade, whereas a clade formed by Chinchilloidea and Octodontoidea is less well-supported (Upham and Patterson,

2012; Voloch et al., 2013). Inter-familial relationships within superfamilies, on the other hand, are much less strongly supported (Upham and Patterson, in press).

- (b) *Sequence and timing of major radiations.* Several major radiations, such as that of spiny rats and allies (Echimyidae, 88 species) and tuco-tucos (Ctenomyidae, 65 species) have been difficult to resolve in terms of phylogenetic relations or even number of species (e.g., Parada et al., 2011). In addition, timing estimates based on a few genes are accompanied by large confidence intervals, making it difficult to tie these events to specific geological and climatic episodes (Opazo, 2005). Similarly, exchanges between regions and biomes and, especially their timing, have been insufficiently resolved with limited genetic data (e.g., Upham et al., 2013).
- (c) *Molecular basis of evolutionary change.* The unique insulins of caviomorphs have been known for five decades, and analyses of DNA sequences have suggested that positive selection was a driving factor in their divergence (Opazo et al., 2005). The role of proinsulin as a growth factor (De Meyts, 2012), coupled with the unusual developmental biology of caviomorph, suggest that insulins may be involved in some of their unusual adaptations. Because insulin is known to affect the expression of at least 150 genes, a systems approach, rather than a gene-by-gene approach, is required (see MacManes et al., in press). Caviomorph rodents possess changes in several other physiological axes that are otherwise well conserved across mammals (Wriston, 1981), but these changes are not fully understood. Genome-based research in that direction will shed light on both caviomorph biology and important mechanisms of mammalian physiology. On a related line, analyses of complete mitochondrial genomes and selected mitochondrial genes (Tomasco and Lessa, 2011, 2014) of subterranean and non-subterranean octodontoid species have suggested adaptive processes in relation to hypoxic environments. A full understanding of these processes requires analysis of whole genomes to develop a complete picture of the changes involved in the origin of the subterranean lifestyle.
- (d) *Evolutionary genomics of domestication.* Guinea pigs provide a fascinating example of the evolution of domestication (in traditional Andean cultures and into modern biomedical sciences). Molecular phylogenetics suggests that *Cavia tschudii* has been the source of domesticated guinea pigs (Spotorno et al., 2004; Dunnum and Salazar-Bravo, 2010). Analysis of the genomic basis and consequences of domestication will be greatly facilitated by the available guinea pig genome and existing phylogenetic and systematic framework.

SIGMODONTINE RODENTS: AGE, ORIGINS, AND DIVERSITY

Sigmodontines (cotton and rice rats, grass, leaf-eared and vesper mice, and allies) are ubiquitous across South America, from sea level to the high Andes, and from evergreen Amazonia to the Atacama, the world's driest desert. The subfamily Sigmodontinae was the center of extended taxonomic debates during the second half of the XXth century over whether to limit the group, as it is currently understood, to include only the complex-penis cricetids that are predominantly South American (e.g., *Sigmodon*, *Oryzomys*, *Akodon*, *Phyllotis*) or to also include the predominantly

North American forms with a simple, rod-like penis, which are currently placed in the cricetid subfamilies Neotominae and Tylomyinae.

Early DNA-based phylogenies (e.g., Engel et al., 1998; Dubois et al., 1999) strongly supported the monophyly of a group composed of the complex-penis Sigmodontinae to the exclusion of the neotomine-peromyscines and tylomyines. Recent analyses, with broader taxonomic coverage (e.g., Steppan et al., 2004; Ventura et al., 2013), supported those earlier studies. DNA sequence data generally supported a North-Central American origin of sigmodontines at a much earlier date (~ 12 Ma; e.g., Parada et al., 2013 see also Schenk et al., 2013; Vilela et al., 2014) than most scholars were willing to accept before molecular studies (reviewed by Reig, 1981).

As currently defined, the subfamily Sigmodontinae includes 86 extant genera and about 400 extant species, of which 85 genera and 381 species inhabit South America (D'Elia and Pardiñas, in press). Sigmodontine genera have been united into various groups, with some formalized as tribes. The number and content of these groups have varied, depending on the weight given to trenchant characters. Phylogenetic analyses based on DNA sequences prompted important changes in the distinction and composition of tribes. Examples of these changes include: (a) the recognition of the Abrotrichini as distinct from Akodontini, with Scapteromyini subsumed under the latter (Smith and Patton, 1999; D'Elia, 2003); (b) significant redefinition of Phyllotini (Salazar-Bravo et al., 2013) and Thomasomyini (D'Elia et al., 2006), and (c) substantive changes within Oryzomyini (Weksler, 2003). Now, after broad taxonomic sampling and phylogenetic analyses, 73 of 85 genera have been consistently allocated to one of nine recognized tribes (D'Elia and Pardiñas, in press; **Table 2**). However, contemporary understanding of sigmodontine diversification is primarily based on variation at only two loci (the mitochondrial cytochrome b and nuclear IRBP genes) which are incongruent at important parts of the radiation. Finally, 12 genera are *incertae sedis*, including at least one clade of tribal rank (Parada et al., 2013; Ventura et al., 2013).

Several issues of sigmodontine systematics will require considerably greater genetic and taxonomic sampling to be resolved. Among them, we highlight the following:

- (a) *Relationships among tribes.* Contents of currently recognized tribes appear to be relatively stable based on two loci, but phylogenetic relationships among tribes remain uncertain, as does the placement of several distinct genera. At present, only the sister relationship of Sigmodontini + Ichthyomyini and a large clade that includes all other sigmodontines (designated as Oryzomyalia by Steppan et al., 2004) are well supported. Within the Oryzomyalia, Abrotrichini and Wiedomyini are sister to each other, as are Phyllotini and the *incertae sedis* genus *Delomys* (e.g., Ventura et al., 2013), although the latter is based solely on the IRBP locus.
- (b) *The tempo of the sigmodontine diversification: hard vs. soft polytomy.* Given limited genetic sampling, it remains unclear whether the lack of resolution seen at the base of Oryzomyalia reflects a hard polytomy that is the result of an explosive radiation or if it is simply due to lack of resolution for the two loci

Table 2 | Classification and diversity (number of species and genera per tribe) of Sigmodontinae in South America (following Patton et al., in press), with a survey of the available taxonomic coverage for the two most commonly used loci in sigmodontine phylogenetic studies, and available and proposed genomic and transcriptomic coverage.

Tribe	Genera (species)	Genes most commonly used in phylogenetics		Available mitochondrial genomes	Available* and proposed (minimum) coverage	
		Mitochondrial cyt b	Nuclear IRBP		Genomes	Transcriptomes
Abrotrichini	5 (13)	5 (12)	5 (11)			1*
Akodontini	15 (85)	13 (63)	13 (35)	1 1		1
Ichthyomyini	4 (13)	0 0	0 0			1
Oryzomyini	30 (118)	28 (78)	29 (63)			1
Phyllotini	11 (50)	11 (43)	11 (22)			1
Reithrodontini	1 (2)	1 (1)	1 (1)			1
Sigmodontini	1 (4)	1 (3)	1 (3)		1	1
Thomasomyini	5 (73)	4 (27)	4 (10)			1
Wiedomyini	1 (2)	1 (2)	1 (1)	1 1		1
<i>incertae sedis</i>	12 (21)	10 (15)	10 (10)			1

Mitochondrial genomes reported in Vilela et al. (2014); transcriptome in Giorello et al. (2014).

so far employed. In either case, adoption of a phylogenomic approach would help clarify this issue.

- (c) **Historical Biogeography.** One of the main aspects of sigmodontine historical biogeography that remains unresolved is the geographic placement of the basal sigmodontine radiation, which in turn relates to the number of sigmodontine lineages that independently invaded South America. To solve this issue and provide a well-resolved phylogeny among sigmodontine tribes, clarification of the placement of Sigmodontinae within Cricetidae (e.g., sister to Tylomyinae as suggested by Parada et al., 2013) is needed. Further, study of sigmodontine relationships with sufficiently dense taxonomic coverage will allow robust inference of the geographic distribution of the sigmodontines most recent common ancestor.

A PHYLOGENOMICS RESEARCH AGENDA FOR SOUTH AMERICAN RODENTS

A concerted effort is needed to propel the study of South American rodents into the genomic era (broadly taken to encompass any large-scale surveys of genetic variation, including transcriptomics). Currently, public databases include genomic data for three caviomorphs (domestic guinea pig, chinchilla, and degu), representing three of the four caviomorph superfamilies. Reported transcriptomes include those three species, plus an additional guinea pig and the social tuco-tuco. There are no genomic/transcriptomic data for 7 families (including the

single one within the superfamily Erethizontoidea) or for the only non-South American family Capromyidae (limited to the West Indies; see Upham and Patterson, in press, for discussion of capromyid rank and phylogenetic position). Currently, there are complete mitochondrial genomes available for 11 caviomorphs (Tomasco and Lessa, 2011; Voloch et al., 2013). Sigmodontines lag far behind in spite of their impressive diversity. To our knowledge, no genomes are available or under way, and the only transcriptomes currently available come from the olive mouse (*Abrothrix olivacea*, Giorello et al., 2014). Only two sigmodontine mitochondrial genomes are currently available (Vilela et al., 2014).

Naturally, genomes offer more data than transcriptomes, but the latter provide ready access to the sequences of thousands of genes at a much smaller cost and are much more easily assembled. At this stage, we propose a mixed strategy to include 2 new genomes, as well as transcriptomes of 16 species, generating the following new data: (a) the genome of an erethizontoid to represent the fourth caviomorph superfamily; (b) transcriptomes of the 7 families of caviomorphs (including the Capromyidae) for which such data are currently unavailable; (c) the genome of one sigmodontine species; and (d) 9 transcriptomes to complete the representation of the ca. 10 recognized clades of tribal rank (Parada et al., 2013) of sigmodontine rodents. The inclusion of additional, divergent lineages of sigmodontines currently not assigned to tribes should be considered, as should

relevant outgroup taxa for both caviomorphs and sigmodontines. Sequence data should be deposited in public databases and referenced to museum voucher specimens and associated data (see below). In addition, we propose generating a specific database to facilitate studies of Neotropical rodents, cross-referenced to both museum and genomic databases. This effort should aim to generate a collection of some 4000 full-length (>95% coverage of codons), annotated and aligned orthologous loci, with the corresponding intron/exon structure referenced to available genomes.

We envision these data to be widely utilized in several ways by students of Neotropical rodents. Relative to systematics: (a) phylogenomic studies can be carried out by the direct generation of transcriptomes and/or genomes of groups of interest (Lindblad-Tohl et al., 2011; Lin et al., 2014); (b) exon capture protocols can capitalize on reference sequences (Bi et al., 2012, 2013; Nachmann, 2013); and (c) primer sets can be developed to target selected sets of genes (e.g., Perelman et al., 2011). But many other studies of specific biological systems (e.g., caviomorph insulin and related genes) will be facilitated by genome-level data.

STANDARDS FOR VOUCHER SPECIMENS: A ROLE FOR NATURAL HISTORY COLLECTIONS

Natural history collections are essential repositories of voucher specimens that ensure reproducible science, traceability and extension of research through access to supplemental data and biological samples associated with vouchers. In recent years, there has been a growing recognition of the importance of cross-references between museum collections and DNA sequence data (Federhen et al., 2009). For example, the Genome 10K Project (genome10k.soe.ucsc.edu) and Barcode of Life (barcodeoflife.org) initiatives have developed specific recommendations in this respect. Increasingly museum databases (e.g., arctos.database.museum) are available on the web and cross-referenced with GenBank and other web-based databases and applications (e.g., GIS tools). Building on these practices, we propose the following standards for vouchers associated with our initiative:

- (a) In phylogenomic studies sampling should be designed to maximize the usage of type or topotypic specimens and type taxa.
- (b) Specimens should be collected by adhering to laws and permits that govern collection and transport.
- (c) Standard documentation of date, georeferenced collection locality, external measurements, sex and associated reproductive data, habitat notes and digital locality photos should be produced and deposited in association with the specimens.
- (d) Series of specimens for each rodent species should be preserved using a variety of standard preparations including standard skeleton + skin and fluid-preserved specimens, with samples of liver, heart, lung, spleen, and kidney immediately frozen in liquid nitrogen (Yates et al., 1996; Wong et al., 2012). Subsamples of tissues should be preserved in buffers (e.g., RNAlater®) that preserve host RNA and RNA viruses (Torres-Pérez et al., 2011; Dacheux et al., 2014). All host

specimens should be screened for ecto and endoparasites, with representative subsets of parasite specimens frozen in liquid nitrogen and the remainder properly prepared and preserved in 95% ethanol.

- (e) Transcriptomes preferably should be based on RNA sequences from multiple organs to maximize gene recovery.
- (f) All specimens should be deposited in established natural history museums accessible to the research community. Associated data and records should be digitized and available via the web, with hosts and parasites cross-referenced to each other, and as genomes are generated, data should be freely available and tied to the specimens and to permanent data repositories such as GenBank and associated resources. The goal is to ensure that web-accessible databases can link geo-referenced specimens and all original and derivative data (e.g., Arctos).

CONCLUDING REMARKS

The goals outlined above will be best achieved by a collaborative framework, involving a consortium of natural history museums and other research institutions, and incorporating curators of the corresponding mammal collections and researchers in South American rodent systematics. Specific efforts should be made to involve investigators from multiple countries in Latin America and elsewhere. Such a collective endeavor would serve to tie together diverse research initiatives ranging from field ecological studies or museum-based survey efforts to phylogenomics, functional genomics, and emerging pathogen discovery. Further, it would serve as an example for how to connect and more fully exploit the booming “big data” initiatives in both GIS-based informatics (Peterson et al., 2011) and genomics, all with powerful implications for addressing questions related to environmental and human health (Atlas et al., 2010). Funding is required to cover field and curatorial efforts, sequencing costs, assembly, annotation and databasing. Those efforts, and the involvement of institutions and researchers, can be developed incrementally from an initial core of committed researchers and students, whose participation we hope to encourage. Given the expected fast growth of genomics (broadly understood), many efforts will be carried out by independent investigators. We hope that the systematically oriented framework of our proposal will contribute data, best practices, and standards for new projects.

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